EXHIBIT 3, Tab 2

FOR USE IN OPIOID-TOLERAN

ENTR CHLY

 ${f B}$ only

WARMING:

Oxycodone hydrochloride extendad-release tablets are an epicid agonist and a Schedule II controlled substance with an abuse flability similar to morphine.

Oxycodone can be abused in a manner similar to other opioid agonists, legal or illicit. This should be considered when prescribing or dispensing oxycodone hydrochloride extended-release tablets in situations where the physician or harmacist is concerned about an increased risk of misuse, abuse, or diversion.

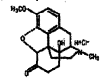
Oxyrodone hydrochleride extended-release tablets are an extended-release orel formulation of oxygotone sydrechloride indicated for the management of mederate to severe paid when a continuous, around-the-clock analysale is needed for an extended period of time.

Daycodons hydrochloride extended-release intilets are NOT intended for use

Expressions Hydrockloride Extended-Release 96 mg Tablets ARE FOR USE 184 creamy TOLERANT PATIENTS DAILY. This tablet strangth may cause fatal respiratory to patients not previously exposed to opioids.

ACOUNT HOROCHLORIDE EXTENDED RELEASE TABLETS. ARE TO SE AWAY SEED WHOLE AND ARE NOT TO SE SONCEN, CHEWED, OR CRUSHED. AND ARE NOT TO SE SONCEN, CHEWED, OR CRUSHED STOROCHLORIDE BY SONCEN, CHEWED, OR CRUSHED STOROCHLORIDE BY DIDED RELEASE TABLETS LEAUS TO RAPID RELEASE AND ASSORPTION OF A POTENTIALLY FATAL DOSE OF OXYCODOME.

Oxycodone Hydrochloride Extended-Release Tablets are an opioid analgesic supplied in 80 mg tablet strength for oral administration. The tablet strength describes the amount of oxycodone per tablet as the hydrochloride sait. The MAR 2.3 ZUL structural formula for exycodone hydrochioride is as follows:



C18H21NO4HCI

M.W. 351.82

The chemical formula is 4,5-Epoxy-14-hydroxy-3-methoxy-17-methylmorphinan-6-one hydrochloride.

Oxycodone is a white, odoriess crystalline powder derived from the opium alkaloid, thebaine. Oxycodone hydrochloride dissolvas in water (t g in 5 to 7 mL). It is slightly soluble in alcohol (octanol water partition coefficient 0.7). Each tablet contains 80 mg of oxycodone hydrochloride. In addition, each tablet contains the following- inactive ingredients: colloidal silicon dioxide. FD&C blue #2 indigo carmine lake, hypromellose (2208, 100M), iron oxide yellow, lactose anhydrous, lactose monohydrate, magnesium stearste, microcrystalline cellulose, polyethylene glycol, titanium dioxide and triacatin.

CLINICAL PHARMACSLOSY

CLINICAL PHARRMACOLORY

Oxycodone is a pure agonist opioid whose principal therapeutic action is analgesia. Other members of the class known as opioid agonists include substances such as morphine, hydromorphone, fentanyl, codeline, and hydrocodons. Pharmacological effects of opioid agonists include anxiotysis, suphroria, feelings of relaxation, respiratory depression, consipation, milosis, and cough suppression, as well as analgesia. Like all pure opioid agonist analgesias, with increasing doses there is increasing analgesia, unlike with tribed agonist/antagonists or non-opioid analgesics, where there is a limit to the analgesic effect with increasing doses. With pure opioid agonist; analgesics, there is no defined maximum dose; the ceiling to analgesic effectiveness is imposed only by side effects, the more serious of which may include somnolence and respiratory depression.

Central Nerveus System

The precise mechanism of the analysis action is unknown. However, specific CNS opioid receptors for endogenous compounds with opioid-like activity have been identified throughout the brain and spinst cord and play a role in the analysis.

Oxycodone produces respiratory depression by direct action on brain stem respiratory centers. The respiratory depression involves both a reduction in the responsiveness of the brain stem respiratory centers to increases in carbon dioxide tension and to electrical stimulation.

Oxycodone degresses the cough reflex by direct affect on the cough center in the medults. Antituseive affects may occur with deets lower than those usually required for analgesia.

Oxycodone causes miosis, even in total darkness. Pinpoint pupils are a sign of opioid overdose but are not pathogrammonic (a.g., pontine lasticus of hemorrhagic or ischemic origin may produce similar findings). Marked mydriasis rather than miosis may be seen with hypoxis in the setting of oxycodone hydrochloride extended-release tablets overdose (See OVERDOSAGE).

Gastroistessissi Tract and Other Smooth Muscle

Oxycodone causes a reduction in motility associated with an increase in smooth muscle tone in the antrum of the stomach and duodenum. Digestion of food in the small intestine is delayed and propulsive contractions are decreased. Propulsive peristaltic waves in the colon are decreased, while tone may be increased to the point of spasm resulting in constipation. Other opinid-induced effects may include a reduction in gastric, bilitary and pancreatic secretions, spasm of sphincies of Oddi, and transient elevations in serum smytase.

Cardiovascular Serioro

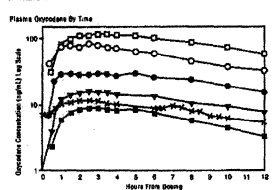
Oxycodone may produce release of histamine with or without associated peripheral vasodilation. Manifestations of histamine release and/or peripheral vasodilation may include pruritus, flushing, red eyes, sweating, and/or orthostatic hypotension.

Gencemiration - Efficacy Relationships Studies in normal volunteers and patients reveal predictable relationships between oxycodone dosage and plasma oxycodone concentrations, as well as between concentration and certain expected optoid effects, such as pupillary constriction, secation, overall "drug affect", analysis and feelings of "retaxation."

As with all opioids, the minimum effective plasma concentration for analysis will vary widely among patients, especially among patients who have been previously

hydrochloride extended-release tablets, steady-state oxycodone from oxyc oxycocone from oxyc.

"hydrochloride scanded-release tablets, seasy-state plattine concentrations to "hydrochloride adended-release tablets. In a study comparing 10 mg of oxycodone hydrochloride extended-release tablets wery 12 hours to 5 mg of immediate-release oxycodone svery 6 hours, the two treatments were found to be equivalent for AUC and Case, and similar for Case, (frough) concentrations. There was less fluctuation in plasma concentrations for the oxycodone hydrochloride extended-release tablets than for the immediate-release formulation.



-14- (i) mg q12h Steady-State

Table 1 Mose (% coefficient variation)					
Regimen	Dosage Form	AUC (ng·hr/mL)†	C _{max} (ng/mL)	T _{max} (hrs)	Trough Cone. (ng/mL)
Single Dose	10 mg oxycodone hydrochloride extended-release tablets	100.7 [26.6]	10.6 [20.1]	2.7 [44.1]	n.a.
	20 mg piyoodone hydrochlorida extended-relates hiblets	207.5 (35.9)	21.4 [36.6]	3.2 [57.9]	ń.a.
	40 mg oxycodone hydrochioride extended-release tablets	423.1 [33.3]	39.3 [34.0]	3.1 [77.4]	n.a.
	80 mg oxycodone hydrochlorida extended-release tablets*	1085.5 {32.3}	88.5 [32.1]	2.1 [52.3]	п.а.
Multiple Dose	10 mg psycodone hydrochloride extended-release tablets q12h	103.6 [38.6]	15.1 (31.0)	3.2 [69.5]	7.2 [48.1]
	5 mg immediate- release q6h	99.0 [36.2]	15.5 {26.8}	1.6 [49.7]	7.4 [50.9]

Tor single-dose AUC-AUC_{D-Int} for multiple-dose AUC-AUC_{D-T}

data obtained while volunteers received nathrexone which can enhance absorption

Table 2 Mean [% coefficient variation)					
Regimen	Dosage Form	AUC (ng-hr/mL)t	C _{max} (ng/mL)	T _{max} (hrs)	Trough Conc. (ng/mL)
Single Doss	4 x 40 mg oxycodone hydrochloride extended-release tablets	1935.3 [34.7]	152.0 {28.9}	256 {42.3}	0,8.
	2 x 80 mg oxycodone hydrochloride extended-release tablets	1859.3 [30.1]	153.4 [25.]]	8\1 [6.63]	8.2.
	1 x 160 mg oxycodone hydrochloride extended-release tablets	1856.4 [36.5]	156.4 [24.8]	254 [36.4]	n.a.

f for single-dose AUC=AUConn; for multiple-dose AUC=AUCo-7
* data obtained while volunteers received natiraxone which can enhance absorption

OXYCODONE HYDROCHLORIDE EXTENDED-RELEASE TABLETS ARE NOT INDICATED FOR RECTAL ADMINISTRATION. Data from a study involving 21 normal volunteen show that conjectione hydrochieride extended-release tablets administered per rectum resulted in an AUC 39% greater and a C_{max} 9% higher than tablet administered by mouth. Therefore, there is an increased risk of adverse event with rectal administration.

Food Effects
Food has no significant effect on the extent of absorption of psycodone from expression flydrochloride antended-release tablets.

apprintered intravenous administration, the volume of distribution (Vss) for oxycodon was 2.6 L/kg. Oxycodone binding to plasme protein at 37°C and a pH of 7.4 wa about 45%. Once absorbed, exycodone is distributed to statetal muscle, live intestinal tract, lungs, splean, and brain. Coycodone has been found in brasst mil

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HYDROCHLONDE EXTENDED-RELEASE TABLETS, 20 mg

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Occodone may produce release of histornine with or without associated peripheral vasodilation. Manifestations of histornine release and/or peripheral vasodilation may include pruritus, flushing, red eyes, sweating, and/or orthostatic hypotension.

Concentration - Efficacy Relationships Studies in normal volunisers and patients reveal predictable relationships between oxycodone dosage and plasma oxycodone concentrations, as well as between concentration and certain expected opioid effects, such as pupillary constriction, sedation, overall "drug effect"; analgesia and feelings of "relaxation."

As with all opioids, the minimum effective plasma concentration for analyseia will vary widely among patients, especially among patients who have been praviously treated with potent agonist opioids. As a result, patients must be treated with individualized litration of dosage to the desired effect. The minimum effective analgesic concentration of psycodone for any individual patient may increase over time due to an increase in pain, the development of a new pain syndrome and/or the development of analgesic tolerance.

Concentration - Adverse Experience Relationships

Conjugation of the Conjugation o clinically relevant.

As with all opinids, the dose must be individualized (see **DOSASE AND ADMINISTRATION**), because the effective analogsic dose for some patients will be too high to be tolerated by other patients.

Pharmacokinetics and Metabelism

The activity of oxygodone hydrochloride extended-release tablets is primarily due to the parent drug oxygodone. Doycodone hydrochloride extended-release tablets are designed to provide extended delivery of oxycodone over 12 hours.

Breaking, chewing or crushing oxycodone invirochloride extended-release lablets eliminates the extended delivery mechanism and results in the rapid release and eliminates the extended delivery mechanism and results in the rapid release and absorption of a potentially fatal dose of oncodone.

Oxycodone release from oxycodone hydrochloride extended-release tablets is pH Oxycodone release from oxycodone hydrochloride extended-release tablets is phindependent. Oxycodone is well absorbed from oxycodone hydrochloride extended-release tablets with an oral bloavallability of 60% to 87%. The relative oral bloavallability of oxycodone hydrochloride extended-release tablets to immediate-release oral dosage forms is 100%. Upon repeated dosing in aormal volunteers in pharmacokinetic studies, steady-state levels were achieved within 24 to 36 hours. Oxse proportionality and/or bloavallability has been established for the 10 mg, 20 mg, 40 mg, 80 mg, and 160 mg tablet strengths for both peek plasma levels (C_{max}) and extent of absorption (AUC). Oxycodone is extensively metabolized and elimination primarily in the urine as both conjugated and innonjugated metabolites. The apparent elimination half-tile of oxycodone following the administration of oxycodone loydrochloride extended-release tablets was 4,5 hours compared to 3,2 hours for immediate release oxycodone. immediate-release oxycodona.

About 50% to 87% of an oral dose of oxycodone reaches the central compartment in comparison to a parenteral dose. This high oral bloavailability is due to low pre-systemic and/or first-pass metabolism. In normal volunteers, the t¹/₂ of absorption is 0.4 hours for immediate-release oral oxycodone. In contrast, oxycodone hydrochloride extended-release tablets, withink a biphasic absorption pattern with two apparent absorption half-times of 0.6 and 5.9 hours, which describes the initial release of oxycodone from the tablet followed by a prolonged release.

Dose proportionality has been established for the 10 mg, 20 mg, 40 mg, 80 mg, and 180 mg tablet strengths for both peak plasma concentrations (C_{max}) and extent of absorption (AUC) (see Table 1 below). Given the short half-life of alimination of

† for single-dose AUC-AUCG-in; for multiple-dose AUC-AUCg-T
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DXYCODONE HYDROCHLONIOE EXTENDED-RELEASE TABLETS ALE NOT INDICATED FOR RECTAL ADMINISTRATION. Data from a study involving 21 normal volunteers show that coycodone hydrochloride extended-release tablets administered per rectum resulted in an AUC 39% greater and a $C_{\rm max}$ 9% higher than tablets administered by mouth. Therefore, there is an increased risk of adverse events with rectal administration.

Food has no significant effect on the extent of absorption of oxycodene from exycodene hydrochloride extended-release tablets.

Following intravenous administration, the volume of distribution (vise) for exycodone was 2.5 L/mg. Oxycodone binding to plasma protein at 37°C and a pH of 7.4 was about 45%. Once absorbed, exycodone is distributed to skeletal muscle, liver, intestinal tract, lungs, spieen, and brain. Oxycodone has been found in breast milk (see PRECAUTIONS).

Occoding hydrochloride is adensively metabolized to noroxycodone, oxymorphone, and their glucuronides. The major circulating matabolite is noroxycodone with an AUC ratio of 6.6 relative to that of oxycodone. Noroxycodone is reported to be a considerably weaker analysis than oxycodone. Oxymorphone, although possessing analysis activity, is present in the plasma only in low consentrations. The correlation between oxymorphone concentrations and opioid effects was much less than that seen with oxycodone plasma concentrations. The analysis activity profile of other metabolites is not known.

The formation of oxymorphone, but not noroxycodone, is mediated by cytochrome P450 206 and, as such; its formation can, in theory, be affected by other drugs (see Drug-Brug intersections).

Conjugations and its metabolites are excreted primarily via the kidney. The amounts measured in the urine have been reported as follows: free cosycodone up to 19%; conjugated cosycodone up to 50%; free cosycodone ON; conjugated cosynorphone 514%; both free and conjugated correspondent have been found in the urine but not quantified. The total plasma clearance was 0.8 L/min for attailts.

Special Populations

The plasma concentrations of oxycodone are only nominally affected by age, being 15% greater in elderly as compared to young subjects.

Fernale subjects have, on average, plasma oxycodone concentrations up to 25% higher than males on a body weight adjusted basis. The reason for this difference

Renal Impairment
Data from a pharmacokinetic study involving 13 patients with mild to severe renal
dysfundsten (creations clearance <60 mL/min) show peak plasma oxycodone and
norcoycodone concentrations 50% and 20% higher, respectively, and AUC values
for oxycodone, norcoxycodone, and oxymorphone 60%, 50%, and 40% higher than
normal subjects, respectively. This is accompanied by an increase in seafation but
not by differences in respiratory rate, pupillary constriction, or several other measures
of drug affect. There was an increase in 11/2 of elimination for oxycodone of only 1 hour
(see PRECAUTIONS).

Data from a study involving 24 patients with mild to moderate hepatic dysfunction show peak plasma expedient and neroxycodone concentrations 50% and 20%. higher respectively, then normal subjects. AUC values are 95% and 65% higher, respectively. Orymorphione peak plaema concentrations and AUC values are lower by 30% and 40%. These differences are accompanied by increases in some, but not other, drug effects, The 1/12 elimination for oxycodone increased by 2.3 hours (see PRESAUTIBRE).

Orug-Brag Sateractions (see PRECAUTIONS)
Oxycodone is membolized in part by cytochrome P450 2DS to oxymorphone which

Case 1:07-cv-07783-Harm 157 or the transmister to ose this rouse of inches to the parties of the process of the parties of the

INDICATIONS AND USAGE

Oxycodone hydrochloride extended-release tablets are an extended-release oral formulation of oxycodone hydrochloride indicated for the management of moderate to severe pain when a continuous, around-the-clock analgesic is needed for an extended period of time.

Oxycodone hydrochloride extended-release tablets are NOT intended for use as a

Physicians should individualize treatment in every case, initiating therapy at the Physicians should enthibute age the earther in every case, introducing hierapy at the appropriate point along a progression from non-opioid analgesics, such as non-steroidal anti-inflammatory drugs and acetaminophen to opioids in a plan of pain management such as outlined by the World Health Organization, the Agency for Health Research and Quelity (formerly known as the Agency for Health Care Policy and Research), the Federation of Stats Medical Boards Model Guidelines, or the American Pain Society.

Oxycodone hydrochloride extended-release tablets are not indicated for pain in the immediate post-operative period (the first 12 to 24 hours following surgery), or if the pain is mild, or not expected to pereist for an extended period of time. Oxycodone hydrochloride extended-release tablets are only indicated for post-operative use if the patient is already receiving the drug prior to surgery or if the post-operative pain is expected to be moderate to severe and persist for an extended period of time. Physicians should individualize treatment, moving from perentarial to oral enablestics as appropriate. (See American Pain Society guidelines.)

CONTRAINDICATIONS

Considerations hydrochioride extended-release tablets are contraindicated in patients with known hypersensitivity to expedient, or in any situation where opioids are contraindicated. This includes patients with significant respiratory depression (in unmonitored settings or the absence of resuscitative equipment), and patients with acute or severe bronchist estima or hypercarbia. Occoding hydrochioride extended-release tablets are contraindicated in any patient who has or is suspected. of having paralytic Heus.

WARNINGS.

DXYCODONE HYDROCHLORIDE EXTENDED-RELEASE TABLETS ARE TO BE DAYCOLDING HYDHOLPHICHEDE EXTENDED-MELEGIS AND TO SE SWALLOWED WHOLE, AND ARE NOT TO BE BROKEN, CHEWED ON CRUSHED. TAKING BROKEN. CHEWED ON CRUSHED OXYCCOONE HYDROCHEGADE EXTENDED-RELEASE TABLETS COULD LEAD TO THE RAPID RELEASE AND ABBORPTION OF A POTENTIALLY FATAL DOSE OF OXYCOOME.

Oxycodone Hydrochloride Extended-Release 40 mg Tablets ARE FOR USE M OPIDIO-TOLERANT PATIENTS ONCY. This limited strongth may cause listed respiratory depression when administered to patients not providusly expended to a judids.

Oxycodess Hydrechloride Extended-Release 56 mg Tablets are for use only in apicid tolerant patients requiring daily coycodese equivalent designs of 168 mg or more. Care should be taken in the prescribing of this tablet strength. Patients should be instructed against ene by individuals other than the patient on whom it was prescribed, as such inappropriate use may have severa medical consequences, including death.

Missee, Abuse and Diversies of Opioids Oxycodone is an opioid agenist of the morphine-type. Such drugs are sought by drug abusers and people with addiction disorders and are subject to criminal diversion.

Oxycodone can be abused in a manner similar to other poloid agonists, legal or illicit. This should be considered when prescribing or dispensing oxycodone hydrochloride extended-release tablets in situations where the physician or pharmacist is concerned. about an increased risk of misuse, abuse, or diversion.

Oxycodone hydrochloride extended-release tablets have been reported as being abused by crushing, chewing, enorting, or injecting the dissolved product. These practices will result in the uncontrolled delivery of the opioid and pose a significant risk to the abuser that could result in overdose and death (see WARNINGS and DAUG ABUSE AND ADDICTION).

Concerns about abuse, addiction, and diversion should not prevent the proper management of pain. The development of addiction to opioid analyseics in properly managed patients with pain has been reported to be rare. However, data are not available to establish the true incidence of addiction in chronic pain patients.

Healthcare professionals should contact their State Professional Licensing Board, or State Controlled Substances Authority for information on how to prevent and detect abuse or diversion of this product.

Interactions with Alcohol and Drugs of Abuse
Oxycodone may be expected to have additive effects when used in conjunction with
alcohol, other opioids, or illicit drugs that cause cantral nervous system depression.

DRUG ABUSE AND ADDICTION

Oxygodica by drochtoriae schaded-release tablets are a me-agenist opicit with an abuse liability similar to morphise and are a Schadule it controlled substance. Oxygodicae, like morphise and other opicids used in analysis, can be abused and is subject to crimical diversion.

Drug addiction is characterized by compulsive use, use for non-medical purposes, and continued use despite harm or risk of harm. Oreg addiction is a treatable disease, utilizing a multi-disciplinary approach, but relapse is common.

"Drug seeking" behavior is very common in addicts and drug abusers. Drug-seeking tactics include amergency calls or visits near the end of office hours, refusal to undergo appropriate examination, testing or referral, repeated "loss" of prescriptions are refusal to provide prior medical records or contact information for other treating physician(s). "Doctor shopping" to obtain additional prescriptions is common among drug abusers and people suffering from

Abuse and addiction are separate and distinct from physical dependence and Abuse and addiction are separate and distinct from physical expensions and tolerance. Physicians should be aware that addiction may not be accompanied by concurrent tolerance and symptoms of physical dependence in all addicts. In addition, abuse of opioids can occur in the absence of true addiction and is characterized by misuse for non-medical purposes, often in combination with other psychoactive substances. Oxygoodone hydrochloride extended-release tablets, like other opioids, have been diverted for non-medical use. Careful record-keeping of prescribing information, including quantity, frequency, and renewal requests is strongly

Proper assessment of the patient, proper prescribing practices, periodic re-evaluation of therapy, and proper dispensing and storage are appropriate measures that help to limit abuse of opioid drugs.

Oxygodone hydrachtoride extended-release tablets are intended for oral use only. Abuse of the crushed tablet poses a hazard of eventees and death. This

The administration of oxygodone may obscure the diagnosis or clinical course in patients with scute abdominal conditions. Oxygodone may aggravate convulsions in patients with convulsive disorders, and all opioids may induce or aggravate seizures in some clinical settings.

setures in some clinical setungs.

Interactions with other CNS Degressents
Chycodone hydrochloride extended-release tablets should be used with caution
and started in a reduced docage (1/8 to 1/2 of the usual docage) in patients who
are concurrently-receiving, other central nervous system degressents including
sedatives of hydrochlor, general anesthetics, phenothazines, other tranquitizers
and alcohot. Interactive effects resulting in respiratory degression, hypotension
profound sedation, or come may result if these drugs are taken in combination with
the usual docas of exycodone hydrochloride extended-release tablets.

interactions with Mixed Agenist/Antagonist Opinic Analysis

Agonist/antigorist analgesics (i.e., pentazodne, natouphine, butorphanol, and bupranorphine) should be administered with caution to a patient who has received or it receiving a course of therapy with a pure opinid agonist analgesic such as oxycodone. In this situation, mixed agonistantagonist analgesic such as natigistic effect of oxycodone and/or may precipitate withdrawal symptoms in

Architectory Surgery and Post-Operative Line
Oxigodesis sydiochloride extended-release tablets are not indicated for pre-empity
analysis (admistalration pre-operatively for the management of post-operative

Expendence hydrochierida extended-release tablets are not indicated for pain in the learnedistic post-operative period (the first 12 to 24 hours following surpery) for patients not proviously taking the crug, because its safety in this selling has not been established.

Oxycodone hydrochieride extended-release tablets are not indicated for pain in the past-operative pariot if the pain is willd or not expected to persist for an extended period of time.

Oxysodome hydrochleride extended-release tablets are only indicated for post-sperative use if the policet in already receiving the drug prior to surgery or if the post-operative gain is expected to be medorate to severe and persist for an extended period of time. Physicians should individualize treatment, moving from parenteral to ural analysaics as appropriate (see American Pain Society

Patients who are already receiving oxycodone hydrochloride extended-release tablets as part of origoing analgesic therapy may be safely continued on the drug if appropriate dosage adjustments are made considering the procedure, other drugs given, and the temporary changes in physiology caused by the surgica intervention (see DOSAGE AND ADMINISTRATION).

Droppidone hydrochloride actanded-release tablets and other morphine-like opi-Large-count myeroconomic excellence resists rathers and other morphise-like op-ords have been shown to decrease bowel motifily. Next is a common post-operative complication, especially after intra-abdominal surgery with optoid analyses. Caution should be taken to monitor for decreased bowel motility in post-operative patients receiving opioids. Standard supportive therapy should be implemented.

Use in Pancrealic/Billary Tract Disease

Oxycodone may cause spasm of the sphinoter of Oddi and should be used with caution in patients with billiary tract disease, including acute pancresitis. Opioids like oxycodone may cause increases in the serum amylase level.

Telerance and Physical Department

Tolerance is the need for increasing doses of opioids to maintain a defined effect such as analogs as (in the absence of disease progression or other external factors). Physical dependence is manifested by withdrawel symptoms after about discontinuation of a drug or upon administration of an antagonist. Physical dependence and tolerance are not unusual during chronic opioid therapy.

The opioid abstinence or withdrawal syndrome is characterized by some or all o the opions assurence of windrawas symptoms is characterized by some of all of the following: residespress, lacrimation, rhinormas, yawning, perspiration, chills myalgia, and mydrasis. Other symptoms also may develop, including: tritability amdely, backache, joint pain, weakness, abdominal cramps, insomnia, nausea angresia, vomiting, diarrhea, or increased blood pressure, respiratory rate, or hear

in general, opioids should not be abruptly discontinued (see DOSAGE ANI ADMINISTRATION: Cassallon of Therapy).

Information for Patients/Caresivers (see PATIENT INFORMATION at the end of the

package insert)
If clinically advisable, patients receiving oxycodone hydrochloride extended-release
tablets or their caregivers should be given the following information by the physician
nurse, pharmacist, or caregiver.

1. Patients should be aware that oxycodone hydrochloride extended-release
tablets contain oxycodone, which is a morphise-like substance.

- Fallents should be advised that oxycodone hydrochloride extended-release tablets were designed to work properly only if swallowed whole. Oxycodon bydrochloride extended-release tablets will release all their contents at once i broken, chewed or enushed, resulting in a risk of tatal overdoes.
- Patients should be advised to report episodes of breakthrough pain an adverse experiences occurring during therapy. Individualization of dosage is essential to make optimal use of this medication.
- Patients should be advised not to adjust the dose of oxycodone hydrochlorid extended-release tablets without consulting the prescribing professional.
- Patients should be advised that population hydrochloride extended-release tablets may impair mental and/or physical ability required for the parformanc of potentially hazardous tables (e.g., driving, operating heavy machinery).
- or pownessy reszeroous tastes (e.g., serving, operating nearly michinary). Patients should not combine oxycodone hydrochlorids extended-releas tablets with alcohol or other central nervous system depressants (steep aids tranquilizers) except by the orders of the prescribing physician, because disagrous additive affects may occur, resulting in serious injury or death. Whomen of childbearing posintial who become, or are planning to become pragnant should be advised to consult their physician regarding the effects a unalgesics and other drug use during pragnancy on themselves and the orders while
- unborn child.
- Patients should be advised that oxycodone hydrochloride extended-releas tablets are a potential drug of abuse. They should protect it from theft, and should move be given to anyone other than the individual for whom it was prescribe
 - Patients should be advised that if they have been receiving treatment wire expectations hydrochloride extended-release tablets for more than a few week and consistent of therapy is indicated, it may be appropriate to taper the expectation of therapy is indicated, it may be appropriate to taper the expectation hydrochloride extended-release tablets dose, rather than abrupt discontinue it, due to the rick of precipitating withdrawal symptoms. The

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an abuse itability similar to manage as a Schedule II controlled substance.

Oxygodone, like merghine as are a Schedule II controlled substance.

2. Patients shrubits were spicials used in analysis, can be abused. and is subject to criminal six

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"Drug seeking" behavior is very common in addicts, and drug abusers. Drug-seeking tactics include emergency calls or visits near the end of office hours, refusal to undergo appropriate examination, testing or referral, repeated "leas" of prescriptions, tampering with prescriptions and reluctance to provide prior medical records or contact information for other treating physician(s). "Doctor shopping" to obtain additional prescriptions is common among drug abusers and people suffering from

Abuse and addiction are separate and distinct from physical dependence and tolerance. Physicians should be aware that addiction may not be accompanied by concurrent tolerance and symptoms of physical dependence in all addicts. In addition, abuse of oploids can occur in the absence of true addiction and is characterized by misuse for non-medical purposes, often in combination with other psychoactive substances. Oxycodone hydrochloride extended-release tablets, like other opioids, have been diverted for non-medical use. Careful record-keeping of prescribing information, including quantity, frequency, and renewal requests is strongly advised.

Proper assessment of the patient, proper prescribing practices, periodic re-evaluation of therapy, and proper dispensing and storage are appropriate measures that help to limit abuse of opioid drugs.

Oversions bytrockloride extended-release tablets are intended for oral use enty. Above of the created tablet posses a fuzzari of overless and death. This risk is increased with concurrent abuse of glockof and other substances. With parasteral abuse, the tablet excludes aim to expected to result in local tissue necrosis, threellow, permonenty gravitoness, and increased risk of escocarditis and valvatar teart injury. Parenteral drug abuse is commonly associated with transmission of infectious diseases such as hepatitis and HV.

Respiratory Degression

Respiratory depression is the chief hazard from oxycodone, the active ingredient in respiratory depression is use care inexate train executions, are every ingrement in convictions in every ingrement in convictions and applied agonitist. Respiratory depression is a particular, problem in eiderly or debitated patients, usually following large initial doses in non-tolerant patients, or when opioids are given in conjunction with other agents that depress respiration.

Oxycodone should be used with extreme caution in patients with significant chronic obstructive pulmonary disease or cor pulmonals, and in patients having a substantially decreased respiratory reserve; hypoxia, hypercapnia, or pre-existing respiratory depression. In such patients, even usual therapeutic doses of oxocorons may decrease respiratory drive to the point of agree, in these patients attempted non-opioid analysics should be considered, and opioids should be employed only under careful medical supervision at the lowest effective dose.

PRESENTATION OF

The respiratory depressant effects of uploids include carbon dioxide retention and secondary elevation of cerebrospinal fluid pressure, and may be markedly exaggerated in the presence of head injury, intracranial lesions, or other sources of pre-existing increased intracranial pressure. Oxygodine produces effects on pupilitary response and consciousness which may obscure neurologic signs of further increases in intracranial pressure in patients with need injuries.

Psyphesistre Effect
Oxycodone hydrochloride extended-release tablets may cause severe hypotension. There is an added risk to individuals whose ability to maintain blood pressure has been compromised by a depleted blood volume, or after concurrent administration with drugs such as phanothistines or other agents which compromise vascenteir tone. Oxycodone may produce orthostatic hypotension in ambulatory patients. Oxycodone, like all opioid analgesics of the morphine-type, should be administered with cardion to patients in circulatory shock, since vascellation produced by the salou may further reduce cardiac output and blood pressure.

PRECAUTIONS

Obioid analgesics have a nerrow therapeutic index in certain patient populations, especially when combined with CRS depressant drugs, and about he reserved for cases where the benefits of opioid analgesia outweigh the known risks of respiratory depression, altered mental state, and postural hypotension.

Use of oxycodone hydrochlorids extended-release tablets is associated with increased potential risks and should be used only with caution in the following conditions: acute alcoholism; adrenocortical insufficiency (e.g., Addison's disease); CNS depression or come; delinium tramens; debilitated patients:

- Patients should be advised that oxycodone hydrochloride extended-release tablets were delight to work properly only if swallowed whole. Oxycodone hydrochloride a derelease tablets will release all their contents at once if broken; chawed or crushed, resulting in a risk of fatal overdose.
- Patients should be advised to report episodes of breakthrough pain and adverse experiences occurring during therapy. Individualization of dosage is essential to make optimal use of this medication.
- Patients should be advised not to adjust the dose of exycodone hydrochloride extended-release tablets without consulting the prescribing professional
- Patients should be advised that oxygodone hydrophioride extended-release tablets may impair mental and/or physical ability required for the performance of gotenitally hazardous tasks (e.g., driving, operating heavy machinery).
- Patients should not combine oxygodone hydrochloride extended-release tablets with alcohol or other central nervous system depressants (steep aids tranquilizers) except by the orders of the prescribing physician, because dangerous additive affects may occur, resulting in serious lajury or death.
- Women of childbearing potantial who become or are planning to become pregnant should be advised to consult their physician regarding the effects of analysiss and other drug use during pregnancy on themselves and their unborn child.
- Patients should be advised that oxycodone hydrochloride extended-release tablets are a potential drug of abuse. They should protect it from theft, and it should rever be given to anyone other than the individual for whom it was prescribed.
- Patients should be advised that if they have been receiving treatment with oxycodone hydrochloride extended-release tablets for more than a few weeks and cossation of therapy is indicated, it may be appropriate to taper the oxycodone hydrochloride extended-release tablets dose, rather than abruptly discontinue it, due to the risk of precipitating withdrawal symptome. Their physician can provide a dose schadule to accomplish a gradual discontinuation of the madication.
- 10. Patients should be instructed to keep oxycodone hydrochloride extended-release tablets in a secure place out of the reach of children. When oxycodone hydrochloride extended-release tablets are no longer needed, the unused tablets should be destroyed by flushing down the tollet.

Use in Drug and Alephol Addiction

Oxycodone hydrochloride extended-release tablets are an oploid with no approved use in the management of addictive disorders. Their proper usage in individuals with drug or alcohol dependence, either active or in remission, is for the management of pain requiring opioid analysis.

Drug-Drug Interactions

Opioid analgestics, lociuding oxycodone hydrochloride extended release tablets, may unhance the neuromuscular blocking action of skeletal muscle relevants and produce an increased degree of respiratory depression.

Oxycodone is matabolized in part to oxymorphone via cytochrome P450 206. While this pathway may be blocked by a variety of drugs (e.g., certain cardiovascular drugs including amiodarone and quinifien as well as polycyclic entidepressants), such blockade has not yet been shown to be of clinical significance with this agent. Clinicals should be aware of the possible interaction, however.

Use with CHS Depressants

Uses with CRS begressions. Drycocone hydrochloride extended-release tablets, like all oploid analysists, should be started at 1/3 to 1/2 of the usual docage in patients who are concurrently receiving other cintral nervous system depressants including sedatives or hypnotics, general anathetics, phenothiazines, centrally acting anti-enetics, tranquilizers, and alcohol because respiratory depression, hypotension, and profound edation or come may result. No specific interaction between oxycodone and monoamine oxidase inhibitors has been observed, but causion in the use of any opioid in patients taking this class of drugs is appropriate.

Carcinogenests, Mutagenesis, languirment of Festility Studies of psycodone to evaluate its carcinogenic potential have not been conducted.

Oxycodone was not mutagenic in the following assays: Ames Salmonella and E. coli Oxycodone was not mutagenic in the following assays: Ames Salmonella and E. coli-test with and without metabolic activation at does of up to 5000 mcg, chromosomal aborration test in human lymphocytes in the absence of metabolic activation at doese of up to 1500 mcg/mL, and with activation 48 hours after exposure at doese of up to 5000 mcg/mL, and in the *in vivo* bone marrow micronucleus test in mice (at plasma levels of up to 48 mcg/mL). Oxycodone was clastogenic in the human lymphocyte chromosomal assay in the presence of metabolic activation in the human chromosomal aborration test (at greater than or equal to 1250 mcg/mL), at 24 but not 48 hours of exposure and in the mouse lymphoma assay at doese of 50 mcg/mL or greater with metabolic activation and at 400 mcg/mL or greater without metabolic activation. 76-14

Toratogenic Effects—Category B. Juction studies have been performed in rats and rabbits by oral administration at coses up to 8 mg/kg and 125 mg/kg, respectively. These doses are 3 and 46 times a human dose of 160 mg/kg, based on mg/kg basis. The results did not reveal evidence of herm to the fetus due to exycodone. There are, however, no adequate and well-controlled studies in pregnant women. Because animal reproduction studies are not always predictive of human response. this drug should be used during pregnancy only if clearly needed.

Latter and Delivery Oxygodone hydrochloride extended-release tablets are not recommended for use exceptions nyarogatoriae extensed-release takens are not recommended for use in women during and immediately prior to labor and delivery because oral oploids may cause respiratory depression in the newborn. Neonates whose mothers have been taking oxycodone chronically may exhibit respiratory depression and/or withdrawat symptoms, either at birth and/or in the nursery.

Hursing Mothers

Low concentrations of oxycodone have been detected in breast milk. Withdrawat symptoms can occur in breast-feeding inflants when maternal administration of an opioid analgesic is stopped. Ordinarily, nursing should not be undertaken while a patient is receiving oxycodone hydrochloride extended-release tablets because of the possibility of sedation and/or respiratory depression in the infant.

Safety and effectiveness of exycodone hydrochloride extended-release tablets have not been established in pediatric patients below the age of 18. It must be remembered that anycogone hydrochloride extended-release tablets cannot be crushed or divided for administration.

Gerlatric Use

in controlled pharmacokinetic studies in elderly subjects (greater than 65 years) the clearance of psycodone appeared to be slightly reduced. Compared to young adults, the plasma concentrations of psycodone were increased approximately 15% (see PHARMACOKINETICS AND METADOLISM). Of the total number of subjects (445) in clinical studies of oxycodons hydrochloride extended-release tablets, 148 (33.3%) were age 55 and older (including those age 75 and older) while 40 (9.0%); were age 75 and older. In clinical trials with appropriate initiation of therapy and loss titration, no unlowerd or unexpected side effects were seen in the elderly patients who received oxycodone hydrochloride extended-release tablets. Thus, the usual doses and dosing intervals are appropriate for these patients. As with all opioids, the starting dose should be reduced to 1/3 to 1/2 of the usual dosage in debilitated, non-tolerant patients. Respiratory depression is the chief fizzard in elderly or debilitated patients, usually following large initial doses in non-tolerant patients. or when opioids are given in conjunction with other agents that depress respiration.

Laboratory Monitoring

Due to the broad range of plasma concentrations seen in clinical populations, the varying degrees of pain, and the development of tolerance, plasma oxycodone measurements are usually not helpful in clinical management. Plasma concentrations of the active drug substance may be of value in selected, unusual or complex cases.

Hopatic impairment

A study of progradione hydrochloride extended-release tablets in patients with nepatic impairment indicates greater plasma concentrations than those with normal function. The initiation of therapy at 1/3 to 1/2 the usual doses and careful dose titration is warranted.

Read Impairment

In patients with renal impairment, as evidenced by decreased creatinine clearance (<60 mL/min), the concentrations of oxycodone in the plasma are approximately 50% higher than in subjects with normal renat function. Dose initiation should follow a conservative approach. Dosages should be adjusted according to the clinical situation.

In pharmacoldinetic studies, optoid-naive females demonstrate up to 25% higher average plasme concentrations and greater frequency of typical optoid adverse events than males, even after adjustment for body weight. The clinical relevance of a difference of this magnitude is low for a drug intended for chronic usage at individualized dosages, and there was no male/female difference detected for afficacy. or adverse events in clinical trials.

ADVERSE REACTIONS

The safety of coycodone hydrochloride adanded-release tablets was evaluated in double-blind clinical trials involving 713 patients with moderate to severe pain of various etiologies; in open-label stadies of cancer pain, 187 patients received oxycodone hydrochloride extended-release tablets in total daily dose a ranging from 20 mg to 640 mg per day. The average total daily dose was approximately 105 mg per day.

Serious adverse reactions which may be associated with oxygodone hydrochloride extended-release tablet therapy in clinical use are those observed with other opioid analgesics, including respiratory depression, apnea, respiratory errest, and (to an even lesser degree) circulatory depression, hypotension, or shock (see OVERDOSAGE).

The pure opioid antage such as naloxone or naimefene are specific antidotes against respiratory de in from opioid overdose. Opteid antagonists should not be administered in the assence of clinically eignificant respiratory or circulatory depression secondary to exycodone overdose. In patients who are physically dependent on any opioid agonist including oxycodone hydrochloride extended-release tablets, an abrupt or complete reversal of opioid effects may precipitate an acute abstinence syndrome. The severity of the withdrawal syndrome produced will depend on the degree of physical dependence and the dose of the antagonist administered. Please see the prescribing information for the specific opioid antagonist for details of their proper use.

DOSAGE AND ADMINISTRATION

GOMEN PRINCIPLES

OXYCODONE HYUNOCHLORIDE EXTENDED-RELEASE TABLETS ARE AN OPIDIO
AGONIST AND A SCHEDULE IL CONTROLLED SUBSTANCE WITH AN ABUSE
LIABILITY SIMILAR TO MORPHINE.

OXYCODONE, LIKE MORPHINE AND OTHER OPIDIDS USED IN ANALGEBIA, CAN BE ABUSED AND IS SUBJECT TO CRIMINAL DIVERSION.

OXYGODONE MYDROCHLORIDE EXTENDED-RELEASE TABLETS ARE TO BE SWALLOWED WHOLE, AND ARE NOT TO BE BROKEN, CHEWED OR CRUSHED. TAKING BROKEN, CHEWED OR CRUSHED OXYGODONE HYDROCHLORIDE EXTENDED-RELEASE AND ABSORPTION OF A POTENTIALLY PATAL DOSE OF OXYGODONE.

in treating pain it is vital to assess the patient regularly and systematically. Therapy should also be regularly reviewed and adjusted based upon the patient's own reports of pain and side effects and the health professional's clinical judgment.

Oxycodone hydrochloride extended-release tablets are indicated for the management of moderate to severe pain requiring treatment with a strong apioid for continuous, around-the-clock analysis for an extended period of time. The extended-release nature of the formulation allows the expectation of hydrochloride extended-release stablets to be effectively administered every 12 hours (see CLINICAL PHARMACOLOGY; PHARMACOL PM) dosing, tailored to their pain pattern. It is usually appropriate to treat a patient with only one opioid for around-the-clock therapy.

Physicians should individualize treatment using a progressive plan of pain management such as outlined by the World Health Organization, the American Pain Society and the Federation of State Medical Boards Model Guidelines. Health care professionals should follow appropriate pain management principles of careful assessment and ongoing monitoring [See BOXED WARNINGS].

Initiation of Therapy
It is critical to initiate the dosing regimen for each patient individually, taking into account the patient's prior opioid and non-opioid analysis treatment. Attention should be given to:

the general condition and medical status of the patient;

- the daily dose, potency and kind of the analgesic(s) the patient has been taking;
- the reliability of the conversion estimate used to calculate the dose of oxygodone:
- the patient's opioid exposure and opioid tolerance (if any);
- special safety issues associated with conversion to oxygodone hydrochloride extended-release tablets doses at or exceeding 160 mg g12h (see Special instructions for Oxygodone Hydrochloride Educated-Release Tablets, 28 mg); and
- the balance between pain control and adverse experiences.

Care should be taken to use low initial doses of expendence hydrochloride extended-release labiets in patients who are not already opioid-tolerant, especially those who are receiving concurrent treatment with muscle relexants, sedatives, or other CNS active medications (see PRECAUTIONS: Drug-Brug interactions).

For initiation of oxycodone hydrochloride extended-release tablets therapy for patients previously taking opioids, the conversion ratios from Foley, KM. [NEJM, 1985; 313;84-95], found below, are a reasonable starting point, although not verified in well-controlled, multiple-dose trials.

Oxycodone hydrochloride extended-release tablets should be individually titrated to a dose that provides adequate analgesia and minimizes side effects.

1. Using standard conversion ratio estimates (see Table 4 below), multiply the

- mg/day of the previous opioids by the appropriate multiplication factors to obtain the equivalent total daily dose of oral exycodone.
- When converting from exycodone, divide this 24-hour exycodone dose in half to obtain the twice a day (q12h) dose of exycodone hydrochloride extended-
- Round down to a dose which is appropriate for the tablet strength available (80 mg tablets).





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OXYCODONE
HYDROCHLORIDE EXTENDI
RELEASE TABLETS,
0033

Casse 11 07/ cov south from a submark or page in which the page in the conversion ratios from Foley, it is individualized dosage, and there we individualized dosages, and there we individualized dosages are supported to the individualized dosages. or adverse events in clinical trials.

ADVERSE REACTIONS

The safety of oxycodone hydrochloride extended-release tablets was evaluated in double-blind clinical trials involving 713 patients with moderate to severe pain of various etiologies. In open-label studies of cancer pain, 187 patients received oxycodone hydrochloride extanded-release tablets in total daily doses ranging from 20 mg to 640 mg per day. The average total daily dose was approximately 105 mg per day.

Serious adverse reactions which may be associated with oxycodone hydrochloride extended-release tablet therapy in clinical use are those observed with other opioid analgesics, including respiratory depression, apnea, respiratory arrest, and (to an even lesser degree) circulatory depression, hypotension, or shock (see OVERDUSAGE).

The non-serious adverse events seen on initiation of therapy with oxygodone lystroulistics oxigned-release takers are typical uploté side effects. These events are dose-dependent, and their frequency depends upon the dose, the clinical setting, the patient's level of opioid-tolerance, and host factors specific to the individual. They should be expected and managed as a part of opioid-analgesia. The most frequent (55%) include: constipation, hauses, somatience, dizziness, vomiting, prunius, headache, dry mouth, sweating, and asthenia.

In many cases the frequency of these events during initiation of therapy may be minimized by careful individualization of starting dosage, slow thration, and the avoidance of large swings in the plasma concentrations of the opioid. Many of these adverse events will case or decrease in intensity as oxygodone hydrochloride extended-release tablets therapy is continued and some degree of tolerance is

Clinical trials comparing exprodone hydrochloride extended-release tablets with immediate-release expressions and placebo, revealed a similar adverse event profile between expressions hydrochloride extended-release tablets and immediate-release expressions. The most common adverse events (>5%) reported by patients at least once during therapy were:

,	Covcodene Hydrochloride Extended-Release Tablets (n=227)	mmediate-Release (n=225)	Placebo (n=45)
	(%)	(%)	(%)
Constinution	23	26 27	7
Nausea	23	27	11
Somnoience	23	24	4
Dizziness -	13	16	9
Pruritus	13	12	2
Vomiting	12	14	7
Headache	7	8	7
Dry Mouth	6	7	2
Asthenia	6	7	
Sweating	5	6	2

The following adverse experiences were reported in oxycodone hydrochloride extended-rolesse tablets treated patients with an incidence between 1% and 5%. In descending order of frequency they were anorsia, nervousness, insomnia, rever, confusion, diarrives, abdominal pain, dyspepsia, rest, artiety, euphoria, dyspres, postural hypotension, chills, twitching, gastritis, abnormal dreams, thought abnormalities, and hiccups.

The following adverse reactions occurred in less than 1% of patients involved in clinical trials or were reported in post marketing experience:

Beneral: accidental injury, chest pain, facial edema, malaise, neck pain, pain

Cardiovescular: migraine, syncope, vasodilation, ST depression

Olgastiva: dysphagia, eructation, flatulance, gastrointestinal disorder, increased appetite, nausea and vomiting, stomatitis, lieus

Hemic and Lymphatic: lymphadenopathy

Metabolic and Mutritional: dehydration, edema, hyponatremia, peripheral edema, syndrome of inappropriate antidiuretic hormone secretion, thirst

Nervous: abnormal gait, agitation, amresia, depersonalization, depression, emotional labitty, hallucination, hyperkinesia, hypesthesia, hypotoma, makaisa, parasthesia, seizures, speech disorder, stupor, timitus, tremor, vertigo, withdrawal syndrome with

Respiratory: cough increased, pharyngitis, voice alteration

Skin: dry skin ministive dermatitis, exticatia Special Senses: abnormal vision, taste perversion

Urogenital: amenormea, decreased libido, dysuria, hematuria, impotence, polyuria. urinary retention, urinatios impaired

OYERDOSASE

Acute overdosage with expendence can be manifested by respiratory depression, somnolence progressing to stupper or come, ekeletal muscle flaceldity, cold and clammy skin, constricted pupils; bradycardis, hypotension, and death.

Deaths due to overdose have been reported with abuse and misuse of oxycodone Deating due to overdose have been reported with autos and misses of dycotonia hydrochloride extended-release tablets, by ingesting, inhalling, or injecting the crushed tablets. Review of case reports has indicated that the risk of latel overdose is further increased when oxycodone hydrochloride extended-release tablets are abused concurrently with alcohol or other CNS depressants, including other oploids.

in the treatment of oxycodone overdosage, primary attention should be given to the re-establishment of a patent airway and institution of assisted or controlled ventilation. Supportive measures (including oxygen and vasopressors) should be employed in the management of circulatory shock and pulmonary edema accompanying overdose as indicated. Cardiac arrest or arrhythmias may require cardiac massage or defibrillation.

- Oxycodone hydrochloride extended-release tablets should be included to a dose that provides adequate analgesia and minimizes side effects.

 1. Using standard conversion ratio estimates (see Table 4 below), a migritary of the previous opioids by the appropriate multiplication obtain the equivalent total daily dose of oral oxycodone.
- When converting from expendent, divide this 24-hour expendent to obtain the twice a day (q12h) dose of expendent hydrochlorida release tablets.
- Round down to a dose which is appropriate for the tablet streng
- Discontinue all other around-the-clock opioid drugs when hydrochloride extended-release tablets therapy is initi
- No fixed conversion ratio is likely to be satisfactory in all patients patients receiving large opioid doses. The recommended dose Table 4 are only a starting point, and close observation and frequi are indicated until patients are stable on the new therapy.

Yahia 4 Multiplication Factors for Converting the Daily Bose of Prior Opinists to the Dully Bose of Orel Oxycodone' (Mg/Day Prior Opinid × Factor = Mg/Day Graf Oxycodone)

	Oral Prior Opioid	Parenteral Pri
Oxycodone	1	
Codeine	0.15	Napona
Hydrocodone	0.9	
Hydromorphone	4	20
Levorphanol	7.5	15
Meperidine	0.1	0.4
Methadone	1.5	3
Morphine	0.5	3

*Te lie used only for conversion to orsi exysolesie. For patients receivin paranteral opioids, a more conservative conversion is warranted. For e high-dose parenteral morphine, use 1.5 instead of 3 as a multiplication in all cases, supplemental analgesia (see below) should be made avail form of a suitable short-acting analgesic.

Oxycodone hydrochioride extended-release tablets can be safely used co with usual doses of non-opioid analgesics and analgesic adjuvants, pristaken to select a proper initial dose (see PRECAUTIONS).

Conversion from Transfermal Fantanyi to Oxycodene Hydrachioride

Eighteen hours following the removal of the transdermal fentanyl patch, hydrochloride extended-release tablets treatment can be initiated. Alt hydrochronous executorities as taken in eachering an or annual. No has been no systematic assessment of such conversion, a conservative dose, approximately 10 mg q12h of coycodone hydrochloride extentablets should be initially substituted for each 25 mg/hr lentary! I patch. The gatient should be followed closely for early titration, as it limited clinical experience with this conversion.

Minarging Expected Optole Adverse Experiences

Most patients receiving optoids, especially those who are optoid-naive, will side effects. Frequently the side effects from oxygodone hydrochlorid release tablets are transient, but may require evaluation and manageme events such as constigation should be anticipated and treated aggre prophylactically with a stimulant laxative and/or stool softener. Patie usually become tolerant to the constipating effects of opioids.

Other opioid-related side effects such as sedation and neusea are usually and often do not persist beyond the first lew days. If nausea persists and is a to the patient, treatment with anti-emetics or other modalities may n symploms and should be considered.

individualization of Bosspe Crice therapy is initiated, pain relief and other opioid effects should be assessed. Patients should be titrated to adequate effect (generally mik with the regular use of no more than two doses of suppremental analy hours). Patients who experience breakthrough pain may require dosage or rescue medication. Because steady-state plasms concentrations are at within 24 to 36 hours, coapse adjustment may be carried out every 1 t is most appropriate to increase the q12n dose, not the dosing frequent no clinical information on dosing intervals shorter than q12h. As a except for the increase from 10 mg to 20 mg q12h, the total daily coyo usually can be increased by 25% to 50% of the current dose at each it

If signs of excessive oploid-related adverse experiences are observed, the may be reduced. If this adjustment leads to inadequate analgesia, a si dose of immediate-release expections may be given. Alternatively, analgesic adjuvants may be employed. Dose adjustments should I obtain an appropriate balance between pain relief and optoid-relati experiences

If significant adverse events occur before the therapeutic goal of mild c achieved, the events should be treated aggressively. Once adverse under control, upward titration should continue to an acceptable level of ;

During periods of changing analgesic requirements, including initi frequent contact is recommended between physician, other member health-care team, the patient and the caregiver/family.

Special instructions for Oxygenione hydrochloride Extended-Release Tale (For use in epicid-tolerant patients only) Oxygenione Hydrochloride Extended-Release 86 mg Tablets are for opicid-jolerant patients requiring daily oxygenione equivalent desage

Casse 11: 0077-cox-006374835-U-AB of more. Care she

or more. Care should be taken in the presentating of this taken strength. Parlants should be instructed again. By individuals other than the patient for whom it was prescribed, as is appropriate use may have severe medical consequences, including death. Filed 1100/1252/2200077

Supplemental Analysais

Most patients given around-the-clock therapy with extended-release opioids may need to have immediate-release medication available for exacerbations of pain or to prevent pain that occure predictably during certain patient activities (incident pain).

Maintenance of Therapy
The Intent of the thration period is to establish a patient-specific q12h dose that will
maintain adequate analgesia with acceptable aide effects for as long as pain relief
is necessary. Should pain recur then the dose can be incrementally increased to
re-establish pain control. The method of therapy adjustment outlined above should
be acceptable to re-establish pain control. be employed to re-establish pain control.

During chronic therapy, especially for non-cencer pain syndromes, the continued need for around-the-clock opicid therapy should be reassessed periodically (e.g., every 6 to 12 months) as appropriete.

Contation of Therapy

When the patient no longer requires therapy with oxycodone hydrochloride extended-release tablets, doses should be tapered gradually to prevent signs and symptoms of withdrawal in the physically dependent patient.

Conversion from Onyegions Hydrochloride Extended-Release Tablets to Parentaral

To avoid overdose, conservative dose conversion ratios should be followed.

SAFETY AND HANDLING

Oxycodone hydrochloride extended-release tablets are solid dosage forms that contain expections which is a controlled substance. Like morphine, expections is controlled substances act.

Oxycodone hydrochloride extended-release tablets have been targeted for theft and diversion by criminals. Healthcare professionals should contact their State Professional Licensing-Board or State Controlled Substances Authority for Infor-mation on how to prevent and detect abuse or diversion of this product.

HOW SUPPLIED

Controlone Hydrochloride Extended-Release Tablets, 80 mg are green, film-coated, oval, convex tablets debossed with "93" on one side and "33" on the other side. They are available in bottles of 100.

Store at controlled room temperature, between 20" and 25"C (68" and 77"F) (see USP). Dispense in a tight, eight-resistant container as defined in the USP, with a childresistant closure (as required).

CAUTION

DEA Order Form Required.

PATIENT INFORMATION

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OXYCODONE HYDROCHLORIDE EXTENDED-RELEASE TABLETS, 88 mg

Read this interrution carefully before yes take anycolone hydrochieride extended-release tablets. Also read the information you get with your reflix. There may be something new. This information does not take the place of talking with your doctor about your medical condition or your treatment. Only you and your doctor can decide if oxycodone hydrochloride extended-release tablets are right for you. Shere the important information in this leaflet with members of your

What is The Mest Important Information & Should Know About Oxycodene Hydrochloride Edunded-Roleage Tablets?

Lise expedienc hydrochloride axionded-release lablets the way your destartable you in.

- lies expendent hydrochloride extended-rulesse tablets only for the condition for which it was prescribed.
- Oxytedone hydrochloride extended-release tablets are not for accasional ("as needed") use.
- Swallow the tablets whole. Do not break, crush, dissolve, or chew them before swallowing. Oxycodone hydrochloride extended-release tablets work properly over 12 hours only when swallowed whole. If a tablet is breisn, crushed, dissolved, or chewed, the antice 12 hour does will be absorbed into your tody all at each. This can be dangerous, causing an everdeen, and possibly death.
- Keep oxycodous hydrochloride extended-release tablets out of the reach of abilidees. Accidental overdose by a child is dangerous and may result in death.
- Prevent their and misuse. Ocycodone hydrochloride extended-release tablets researt their land initiative that can be a target for people who abuse preschilden inedicines. Therefore, Isbap your tablets in a secure place, to protect them from their. Never give them to amone else. Selling or giving away this medicine is dangerous and against the law.

What are Oxycodone Stylfrochieride Extended-Release Tabletz? Oxycodone hydrochloride extended-release tablets come in several strengths and contain the medicine oxycodone (ox-e-KOE-done). This medicine is a painkillar like morphine. Oxycodone hydrochloride extended-release tablets treat moderate to severe pain that it expected to leaf for an extended period of time. Use oxycodone hydrochloride extended-release tablets regularly during treatment. They contain enough medicine to last for up to twelve hours.

- Who Should Not Take Onycodone Hydrochlaride Extended-Release Tablets?

 Be not take exceedence hydrochlaride extended-release tablets if

 your doctor did not practibe coycodone hydrochlaride extended release tablets for you.
- your pain is mild or will go away in a few days.
- your pain can be controlled by occasional use of other paintillers.
- you have severe asthma or severe lung problems.
- you have had a severe altergic reaction to codeline, hydrocodone, thhydrocodeline, or oxycodone (such as "ylox", Tylanof with Codeline", or Vicodine"). A severe altergic reaction includes a severe rash, hives, breathing problems, or dizziness.
- you had surgery less than 12 to 24 hours ago and you were not taking oxycodone hydrochloride extended-release tablets just before surgery.

Your doctor should know about all your madical conditions before deciding if a trouble breaking or lunk profilems

7 Parce 9 off 110 -id I Avelo Marie Taking Expendence Hydrochloride Ext

- He net drive, c. ... heavy machinery, or participate in any dangerous activities until you know how you react to the company of the company o
- tie not drink zicznoi wiele wing exycodeną bydrochieride az tableta. It may increase the chance of getting dangerous sto
- No not take other medicines without your dector's approval. Include prescription and non-prescription medicines, vitamins, a lie especially careful about products that make you sleepy.

What are the Possiste Side Effects of Oxycodene Hydrochicride Ex Tableta?

Call your dector or get medical help right away it your breathing slows down

you feel faint, dizzy, confused, or have any other unusual sym

Some of the common side effects of oxycodone hydrochloride a tablets are nausez, vomiting, dizziness, drowelness, constigation, itol sweating, weakiness, and headache. Some of these side affects ma continued use.

There is a risk of abuse or addiction with narcotic painkillers. If y drugs in the past, you may have a higher chance of developing abuse again, while using oxycodone hydrochloride extended-release tablishow how often patients with continuing (chronic) pain become addictions. but the risk has been reported to be small

These are not all the possible side effects of oxycodone hydrochloride tablets. For a complete list, ask your doctor or pharmacket.

General Advise About Oxygodene Hydrochloride Extended-Relea • Do not use oxygodene hydrochloride extended-ralease tablet for which it was not prescribed.

- Do not give oxycodone hydrochloride extended-release tablets even if they have the same symptoms you have. Sharing is cause severe medical problems, including death.
- Store oxycodone hydrochloride extended release tablets at I temperature, between 20° and 25°C (68° and 77°F) (see USP

This leaflet summarizes the most important information ab hydrochloride extended-release tablets. If you would like more i with your doctor. Also, you can ask your pharmacist or doctor about coycodone hydrochloride extended-release tablets that is w professionals.

*Tylex and Tylenel with Codelne are brand names of O PHARMAGEUTICAL.

Vicodin is a brand name of ABBOTT LABORATORIES.

Manufactured By: TEVA PHARMAGEUTICALS USA Seliersville, PA 18960

Corcodons hydrochloride extended release tablets come in several strengths and contain the medicine oxycodone (ox-e-KOE-done). This medicine is a paintifier like morphine. Oxycodone hydrochloride extended-release tablets trent moderate to severe pain that is expected to last for an extended period of time. Use oxycodone hydrochloride extended-release tablets regularly during treatment: They contain enough medicine to last for up to twelve hours.

- Who Should Not Take Oxycodone Hydrochloride Extended-Release Tablets?

 De not take oxycodone hydrochloride extended-release tablets if
 your doctor did not prescribe oxycodone hydrochloride extended-release tablets for you.
- your pain is mild or will go away in a few days.
- your pain can be controlled by occasional use of other painkillers.
- YOU have severe astirms or severe king problems.
- you have had a severe alterpic reaction to codeline, hydrococione, dihydrococione, or oxycocione (such as Tylor", Tyleniol with Codeline", or Vicodin""). A severe alterpic reaction includes a severe rash, hives, breathing problems, or dizziness.
- you had surgery less than 12 to 24 hours ago and you were not taking oxycodone hydrochloride extended-release tablets just before surgery.

Your doctor should know about all your medical canditions before deciding if oxycodone hydrochloride extended-release tablets are right for you and what dose is best. Tell your doctor about all of your medical problems, especially the ones listed below:

- trouble breaking or lung problems
- head injury
- liver or iddney problems
- acrenal gland problems, such as Addison's disease
- convulsions or saizures
- alcoholism
- hallucinations or other severe mental problems
- past or present substance abuse or drug addiction

If any of these conditions apply to you, and you haven't told your doctor, then you should tell your doctor before taking oxycodone hydrochloride extended-release

If you are prepared or plan to become prepared, talk with your doctor. Oxycodone hydrophloride extended-release tablets may not be right for you. Tell your doctor if you are breast leading. Oxycodone hydrophloride extended-release tablets will pass through the milk and may harm the baby.

Tell your dector about all the medicines you take, including prescription and non-prescription medicines, vitamins, and kerbal supplements. They may cause serious medical problems when taken with expectance hydrockloride extended-release tablets, especially if they cause drowniness.

- How Should I Take Oxycodone Hydrophioride Extended-Release Tableta?

 Follow your doctor's directions exactly. Your doctor may change your dose based on your reactions to the medicine. Do not change your doctor talls you to change it. Do not take expections hydrochloride extended-release tablets more often than prescribed.
- Swallow the tablets whole. Do not brack, crush, dissolve, or show before swallewing. If the tablets are not whole, your bedy will absorb too much medicing at one time. This can lead to perfous problems, including everdues and death.
- If you make a dose, take it as soon as possible. If it is almost time for your next dose, skip the missed dose and go back to your regular dosing schedule. Do not take 2 doses at once unless your doctor fells you to.
- te case of overlose, call your local emergency number or poison control center right away.
- Review your pain regalarly with your declar to determine if you still need oxygodone hydrochloride extended release tablets.

if you continue to have pale or hothersome side affects, sail your dester.

Stopping exycedane hydrochloride axtended-release tablets. Consult your doctor for instructions on how to stop this medicine slowly to avoid uncomfortable symptoms. You should not stop taking oxycodone hydrochloride extended-release tablets all at once if you have been taking it for more than a few days.

After you stop teking oxygodone hydrochloride axionded-release tablets, flush the unused tablets down the foliet: